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(57) Abstract		
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TITLE METHODS OF PREPARING SOLUBLE, OLIGOMERIC PROTEINS

TECHNICAL FIELD OF THE INVENTION

The present invention relates to a method of preparing soluble oligomeric proteins using recombinant DNA technology.

BACKGROUND OF THE INVENTION

The biological activity of proteins is dependent upon proper tertiary and quaternary structure, or conformation. Many proteins exists as oligomers (structures comprised of two or more polypeptide chains) in their native form. Such oligomers are often stabilized by non-covalent interactions, and are thus dependent on proper tertiary structure of the individual peptides. Expression of a recombinant protein in biologically active form, exhibiting the proper tertiary and quaternary structure, by host cells which do not normally express a native form of the protein, frequently presents a significant challenge. Of particular interest in recombinant protein technology is expression of proteins that are membrane-bound in the biologically active form, as soluble proteins. Soluble proteins are useful as therapeutic agents, and in other applications requiring large quantities of highly purified proteins.

Soluble forms of transmembrane proteins have been prepared by deleting the transmembrane and intracytoplasmic domains, and adding an appropriate signal peptide to enable secretion of the soluble form of the protein (Smith et al., Science 238:1704, 1987; Treiger et al., J. Immunol. 136:4099, 1986). Some soluble proteins have been expressed as fusion proteins in which the extracellular domain of the membrane protein is joined to an immunoglobulin heavy chain constant region (Fanslow et al., J. Immunol. 149:65, 1992; Noelle et al., Proc. Natl. Acad. Sci. U.S.A. 89:6550, 1992), or with the extracellular domain of the murine T lymphocyte antigen CD8 (Hollenbaugh et al., EMBO J. 11:4313, 1992). However, such soluble proteins may not be biologically active due to improper tertiary and/or quaternary structure. Some soluble forms of transmembrane proteins may be biologically active, but poorly expressed, or unstable under the conditions of expression or purification, due to changes in structure as a result of deletion of a portion or portions of the protein.

Leucine zipper is a term that is used to refer to a repetitive heptad motif containing four to five leucine residues present as a conserved domain in several proteins. Leucine zippers fold as short, parallel coiled coils, and are believed to be responsible for oligomerization of the proteins of which they form a domain. Sequences derived from the for and jun leucine zippers have been used in the formation

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of bispecific antibodies by expression of DNA encoding the V_L and V_H regions of antibodies as fusion proteins with the leucine zipper sequences. (Kostelny et al., *J. Immunol.* 148:1547, 1992) Leucine zipper sequences have also been used to replace the dimerization domain of λ repressor, a soluble DNA-binding protein of bacteriophage λ (Hu et al., *Science* 250:1400, 1990), and in the preparation of a dimeric form of MalE, a maltose binding protein of *E. coli* that is exported into the periplasmic space (Blondel and Bedoulle, *Protein Engineering* 4:457, 1991).

There is a need in the art to develop methods of expressing biologically active, recombinant, oligomeric proteins, particularly soluble proteins that are membrane-bound in their biologically active configuration.

SUMMARY OF THE INVENTION

The present invention relates to a method of preparing a soluble, oligomeric mammalian protein by culturing a host cell transformed or transfected with an expression vector encoding a fusion protein comprising a leucine zipper domain and a heterologous mammalian protein. In one embodiment, the heterologous mammalian protein comprises an extracellular domain of a mammalian transmembrane protein; the resulting fusion protein forms an oligomer. In another embodiment, the heterologous mammalian protein comprises a soluble protein such as a cytokine; the resulting fusion protein forms an oligomer. In another embodiment, the leucine zipper domain is removed from the fusion protein, by cleavage with a specific proteolytic enzyme. In another embodiment, a hetero-oligomeric protein is prepared by utilizing leucine zipper domains that preferentially form hetero-oligomers.

BRIEF DESCRIPTION OF THE DRAWINGS

Figure 1A illustrates the ability of soluble, oligomeric human CD40-L comprising a leucine zipper domain to stimulate the proliferation of human tonsillar B cells; Figure 1B illustrates the ability of soluble, oligomeric human CD40-L comprising a leucine zipper domain to stimulate the proliferation of human peripheral blood B cells.

Figure 2 illustrates the inhibition of binding of CD27.Fc to MP.1 cells, which express CD27-L, by a soluble form of CD27-L, sCD27L-3, that comprises a leucine zipper domain.

DETAILED DESCRIPTION OF THE INVENTION

The present invention relates to a method of preparing a soluble mammalian protein by culturing a host cell transformed or transfected with an expression vector encoding a fusion protein comprising a leucine zipper domain and a heterologous mammalian protein. In one embodiment, the heterologous mammalian protein

comprises an extracellular domain of a mammalian transmembrane protein. Exemplary mammalian transmembrane proteins include members of the tumor necrosis factor/nerve growth factor receptor (TNFR/NGFR) family (Farrah and Smith, Nature 358:26, 1992; Goodwin et al., Cell 73:447; 1993), which includes CD40 Ligand (CD40-L), CD27 Ligand (CD27-L), OX40 Ligand (OX40-L), and TNF. Structural studies of certain members of this family of proteins indicate that they form homotrimers. The inventive method will also be useful for other members of this family.

Additionally, many other mammalian transmembrane proteins form oligomers, either hetero-oligomers or homo-oligomers, in their biologically-active form. Members of the hematopoietin receptor family (Cosman et al., Trends Biochem. Sci. 15:265; 1990) are exemplary of such proteins. Gearing et al. (Science 255:1434, 1992) reported the cloning of a gene encoding a protein (gp130) that conferred high-affinity binding to both leukemia-inhibitory factor (LIF) and Oncostatin M (OSM) when expressed in cells along with a low-affinity LIF receptor. Similar interactions of a low-affinity receptor and a second subunit protein, resulting in a high-affinity receptor have also been proposed for other members of this family (Hayashida et al., Proc. Natl. Acad. Sci. U.S.A. 87:0655, 1990; Kitamura et al., Cell 66:1165, 1991; Tavernier et al., Cell 66:1175, 1991; Devos et al., EMBO J. 10:2133, 1991). Soluble forms of the members of the hematopoietin receptor family will exhibit higher affinity for their cognate ligand when expressed as hetero-oliogmers, or in some cases, as homo-oligomers. The same will be true for other transmembrane proteins that comprise two or more subunits.

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In another embodiment, the heterologous mammalian protein comprises a soluble protein such as a cytokine; the resulting fusion protein forms an oligomer. Cytokines are soluble mediators released by cells during an immune or inflammatory response, which provide antigenically non-specific, intracellular signals that are crucial in regulating physiological processes. TNF α , TNF β and certain neurotrophins such as nerve growth factor (NGF) belong to the TNF/NGF family. Modeling studies of certain members of this family indicate that they are likely to form oligomers (Goh and Porter, *Protein Eng.* 4:385, 1991; Peitsch and Jongeneel, *Int. Immunol.* 5:233, 1993). Furthermore, other cytokines, including macrophage colony stimulating factor (M-CSF; Pandit et al., *Science* 258:1358, 1992) are also known to be oligomeric. Such cytokines will also be useful in the inventive method, wherein a leucine zipper domain stabilizes the proper quaternary structure of the oligomeric cytokine.

In another embodiment, hetero-oligomeric forms of cytokines are prepared. A fusion protein of granulocyte-macrophage colony stimulating factor (GM-CSF) and Interleukin-3 (IL-3) has been shown to be a more potent proliferation stimulus than

either factor alone or IL-3 and GM-CSF combined (U.S. Patents 5,073,627 and 5,108,910). Fusion proteins comprising GM-CSF and IL-3 and DNA sequences encoding such fusion proteins are described in U.S. Patents 5,073,627 and 5,108,910, respectively, both of which are incorporated by reference herein. A similar, bivalent protein composed of GM-CSF and IL3 may be formed by the expression of these cytokines as fusion proteins comprising leucine zipper domains that preferentially form heterodimers.

In another embodiment, the leucine zipper domain is removed from the fusion protein, for example by cleavage with a specific proteolytic enzyme. In addition to a leucine zipper sequence and a heterologous protein, such fusion proteins also comprise an amino acid sequence recognized, and cleaved, by a selected proteolytic enzyme. The leucine zipper domain functions to stabilize the recombinant fusion protein during expression and secretion. After purification of the secreted protein, the leucine zipper is enzymatically removed by treating with the proteolytic enzyme. The heterologous protein may then become monomeric. Such monomeric forms of soluble proteins will be useful as receptor antagonists, for example, by binding to a cognate receptor and preventing signaling by preventing cross-linking of the receptor.

Leucine zipper domains

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Leucine zippers were originally identified in several DNA-binding proteins (Landschulz et al., Science 240:1759, 1988). Leucine zipper domain is a term used to refer to a conserved peptide domain present in these (and other) proteins, which is responsible for dimerization of the proteins. The leucine zipper domain (also referred to herein as an oligomerizing, or oligomer-forming, domain) comprises a repetitive heptad repeat, with four or five leucine residues interspersed with other amino acids. Examples of leucine zipper domains are those found in the yeast transcription factor GCN4 and a heat-stable DNA-binding protein found in rat liver (C/EBP, Landschulz et al., Science 243:1681, 1989). Two nuclear transforming proteins, fos and jun, also exhibit leucine zipper domains, as does the gene product of the marine proto-oncogene, c-myc (Landschulz et al., Science 240:1759, 1988). The products of the nuclear oncogenes fos and jun comprise leucine zipper domains preferentially form a heterodimer (O'Shea et al., Science 245:646, 1989; Turner and Tjian, Science 243:1689, 1989). The leucine zipper domain is necessary for biological activity (DNA binding) in these proteins.

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The fusogenic proteins of several different viruses, including paramyxovirus, coronavirus, measles virus and many retroviruses, also possess leucine zipper domains (Buckland and Wild, Nature 338:547,1989; Britton, Nature 353:394, 1991; Delwart and Mosialos, AIDS Research and Human Retroviruses 6:703, 1990). The leucine

zipper domains in these fusogenic viral proteins are near the transmembrane region of the proteins; it has been suggested that the leucine zipper domains could contribute to the oligomeric structure of the fusogenic proteins. Oligomerization of fusogenic viral proteins is involved in fusion pore formation (Spruce et al. *Proc. Natl. Acad. Sci. U.S.A.* 88:3523, 1991). Leucine zipper domains have also been recently reported to play a role in oligomerization of heat-shock transcription factors (Rabindran et al., *Science* 259:230, 1993).

Leucine zipper domains fold as short, parallel coiled coils. (O'Shea et al., Science 254:539; 1991) The general architecture of the parallel coiled coil has been well characterized, with a "knobs-into-holes" packing as proposed by Crick in 1953 (Acta Crystallogr. 6:689). The dimer formed by a leucine zipper domain is stabilized by the heptad repeat, designated (abcdefg)_n according to the notation of McLachlan and Stewart (J. Mol. Biol. 98:293; 1975), in which residues a and d are generally hydrophobic residues, with d being a leucine, which line up on the same face of a helix. Oppositely-charged residues commonly occur at positions g and e. Thus, in a parallel coiled coil formed from two helical leucine zipper domains, the "knobs" formed by the hydrophobic side chains of the first helix are packed into the "holes" formed between the side chains of the second helix.

The leucine residues at position d contribute large hydrophobic stabilization energies, and are important for dimer formation (Krystek et al., Int. J. Peptide Res. 38:229, 1991). Lovejoy et al. recently reported the synthesis of a triple-stranded α-helical bundle in which the helices run up-up-down (Science 259:1288, 1993). Their studies confirmed that hydrophobic stabilization energy provides the main driving force for the formation of coiled coils from helical monomers. These studies also indicate that electrostatic interactions contribute to the stoichiometry and geometry of coiled coils.

Several studies have indicated that conservative amino acids may be substituted for individual leucine residues with minimal decrease in the ability to dimerize; multiple changes, however, usually result in loss of this ability (Landschulz et al., Science 243:1681, 1989; Turner and Tjian, Science 243:1689, 1989; Hu et al., Science 250:1400, 1990). van Heekeren et al. reported that a number of different amino residues can be substituted for the leucine residues in the leucine zipper domain of GCN4, and further found that some GCN4 proteins containing two ieucine substitutions were weakly active (Nucl. Acids Res. 20:3721, 1992). Mutation of the first and second heptadic leucines of the leucine zipper domain of the measles virus fusion protein (MVF) did not affect syncytium formation (a measure of virally-induced cell fusion); however, mutation of all four leucine residues prevented fusion completely

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(Buckland et al., J. Gen. Virol. 73:1703, 1992). None of the mutations affected the ability of MVF to form a tetramer.

Recently, amino acid substitutions in the a and d residues of a synthetic peptide representing the GCN4 leucine zipper domain have been found to change the oligomerization properties of the leucine zipper domain (Alber, Sixth Symposium of the Protein Society, San Diego, CA). When all residues at position a are changed to isoleucine, the leucine zipper still forms a parallel dimer. When, in addition to this change, all leucine residues at position d are also changed to isoleucine, the resultant peptide spontaneously forms a trimeric parallel coiled coil in solution. Substituting all amino acids at position d with isoleucine and at position a with leucine results in a peptide that tetramerizes. Peptides containing these substitutions are still referred to as leucine zipper domains since the mechanism of oligomer formation is believed to be the same as that for traditional leucine zipper domains such as those described above. However, prior to the present invention, the effect of these substitutions upon longer peptides of which the leucine zipper is but a small domain was not known, nor was it known if peptides comprising these sequences could be expressed and secreted by cells.

Preparation of Gene Fragments and Oligonucleotides

Oligonucleotide fragments of about 12 to about 20 nucleotides may be prepared according to methods that are known in the art, for example, by using an automated DNA synthesizer. Several such fragments may be synthesized, which encode overlapping portions of a peptide, for example, a leucine zipper domain. Due to the degeneracy of the genetic code, most amino acids are encoded by two or more different nucleotide triplets. The selection of a triplet to encode a given amino acid will depend upon the organism in which the final gene product is to be expressed, among other considerations. Overlapping fragments may then be joined to form a DNA encoding a peptide of interest.

A polymerase chain reaction (PCR) technique (Saiki et al., Science 239:487, 1988) may be employed to amplify gene fragments encoding all or a portion of a protein of interest, using 5' (upstream) and 3' (downstream) oligonucleotide primers derived from the known DNA sequence of the gene, or a gene encoding a related protein. An exemplary set of PCR conditions includes: one cycle at 94°C for 2 minutes, followed by 42°C for two minutes; 30 cycles at 72°C for 1.5 minutes, followed by 94°C for one minute, then 48°C for 1 minute; and one cycle at 72°C for seven minutes. Restriction enzyme sites can also be added to the DNA sequences of interest, in order to facilitate ligation of the resulting PCR product with a plasmid or vector, or with an additional DNA sequence or sequences. Amplified DNA sequences

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may be joined substantially as described by Yon and Fried (Nucleic Acids Res. 17:4895; 1989).

For example, as disclosed in U.S.S.N. 08/097,827, filed July 23, 1993, the disclosure of which is incorporated by reference herein, full length mouse OX40 was cloned using 5' (upstream) and 3' (downstream) oligonucleotide primers based on the published sequence of rat OX40. The upstream primer comprised a recognition site for the restriction endonuclease Spe I upstream of a sequence encoding the first six (N-terminal) amino acids of rat OX40. The downstream primer comprised a recognition site for the restriction endonuclease Spe I upstream of a sequence encoding the last five (C-terminal) amino acids of full-length OX40. The PCR product was digested with Spe I, and an approximately 800 bp fragment was isolated by gel filtration, and used in a second round of PCR reaction. The isolated fragment was ligated into Spe I cut plasmid, pBLUESCRIPT SK® (Stratagene Cloning Systems, La Jolla, CA), which had been treated with calf intestine alkaline phosphatase (CIAP) to prevent self-ligation.

In another example, a DNA encoding only the extracellular region of a transmembrane protein can be obtained by deleting DNA encoding the intracellular and transmembrane portions of the transmembrane protein. Methods to determine which residues should be deleted and for performing the actual deletions are well known in the art. For example, Smith et al. describe a soluble form of the human CD4 antigen prepared by deleting the transmembrane and intracellular portions of the CD4 antigen (Science 238:1704, 1987). Treiger et al. prepared a soluble form of an Interleukin-2 receptor using similar methods using similar methods (J. Immunol. 136:4099, 1986).

A fusion protein may be formed from an extracellular region and a protein (or portion thereof) that is known to be secreted. For example, soluble proteins comprising an extracellular domain from a membrane-bound protein and an immunoglobulin heavy chain constant region was described by Fanslow et al., J. Immunol. 149:65, 1992 and by Noelle et al., Proc. Natl. Acad. Sci. U.S.A. 89:6550, 1992. The extracellular domain of the murine T lymphocyte antigen CD8 has also be utilized to form soluble fusion proteins (Hollenbaugh et al., EMBO J. 11:4313, 1992).

Preparation of Fusion Proteins

Fusion proteins are polypepiides that comprise two or more regions derived from different, or heterologous, proteins or peptides. Fusion proteins are prepared using conventional techniques of enzyme cutting and ligation of fragments from desired sequences. PCR techniques employing synthetic oligonucleotides may be used to prepare and/or amplify the desired fragments. Overlapping synthetic oligonucleotides representing the desired sequences can also be used to prepare DNA constructs encoding fusion proteins. Fusion proteins can comprise several sequences, including a

leader (or signal peptide) sequence, linker sequence, a leucine zipper sequence, or other oligomer-forming sequences, and sequences encoding highly antigenic moieties that provide a means for facile purification or rapid detection of a fusion protein.

Signal peptides facilitate secretion of proteins from cells. An exemplary signal peptide is the amino terminal 25 amino acids of the leader sequence of murine interleukin-7 (IL-7; Namen et al., Nature 333:571; 1988). Other signal peptides may also be employed furthermore, certain nucleotides in the IL-7 leader sequence can be altered without altering the amino acid sequence. Additionally, amino acid changes that do not affect the ability of the IL-7 sequence to act as a leader sequence can be made.

The Flag® octapeptide (SEQ ID NO:1; Hopp et al., BioTechnology 6:1204, 1988) does not alter the biological activity of fusion proteins, is highly antigenic and provides an epitope reversibly bound by a specific monoclonal antibody, enabling rapid detection and facile purification of the expressed fusion protein. The Flag® sequence is also specifically cleaved by bovine mucosal enterokinase at the residue immediately following the Asp-Lys pairing. Fusion proteins capped with this peptide may also be resistant to intracellular degradation in E. coli. A murine monoclonal antibody that binds the Flag® sequence has been deposited with the ATCC under accession number HB 9259; methods of using the antibody in purification of fusion proteins comprising the Flag® sequence are described in U.S. Patent 5,011,912, which is incorporated by reference herein.

· A protein of interest may be linked directly to another protein to form a fusion protein; alternatively, the proteins may be separated by a distance sufficient to ensure that the proteins form proper secondary and tertiary structures. Suitable linker sequences (1) will adopt a flexible extended conformation, (2) will not exhibit a propensity for developing an ordered secondary structure which could interact with the functional domains of fusion proteins, and (3) will have minimal hydrophobic or charged character which could promote interaction with the functional protein domains. Typical surface amino acids in flexible protein regions include Gly, Asn and Ser. Virtually any permutation of amino acid sequences containing Gly, Asn and Ser would be expected to satisfy the above criteria for a linker sequence. Other near neutral amino acids, such as Thr and Ala, may also be used in the linker sequence. The length of the linker sequence may vary without significantly affecting the biological activity of the fusion protein. Linker sequences are unnecessary where the proteins being fused have non-essential N- or C-terminal amino acid regions which can be used to separate the functional domains and prevent steric interference. Exemplary linker sequences are described in U.S. patents 5,073,627 and 5,108,910, the disclosures of which are incorporated by reference herein.

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When an oligomeric fusion protein is formed from the extracellular portion of a transmembrane protein, a DNA sequence encoding an oligomer-forming domain, such as a leucine zipper domain, is fused to a DNA sequence encoding the extracellular region of the transmembrane protein. The members of the fusion protein are joined such that the oligomer-forming domain of the fusion protein is located in the same orientation relative to the fusion protein as the transmembrane and intracytoplasmic reigns of the native transmembrane protein. An oligomeric fusion protein will be stabilized by the coiled-coil interaction of leucine zipper domain. Thus, in one example, a fusion protein comprising an extracellular region derived from a ligand for CD40 (CD40-L), a type II transmembrane protein described in U.S.S.N. 07/969,703. the disclosure of which is incorporated by reference herein, the oligomer-forming domain, a leucine zipper sequence, is fused to the amino-proximal end of the extracellular region. In a fusion protein derived from a type I transmembrane protein, the oligomer-forming domain would be fused to the carboxy-proximal end of the extracellular region of the type I transmembrane protein. Other transmembrane proteins traverse the cell membrane more than once. Such transmembrane proteins will have two or more different extracellular regions. Soluble, oligomeric fusion proteins may also be prepared from two or more of such different extracellular regions from the same transmembrane protein.

Oligomeric forms of proteins that occur naturally in soluble form may also be prepared. In such cases, the oligomer-forming domain is joined to the soluble protein such that formation of an oligomer follows the conformation of the biologically active, soluble protein. Furthermore, either homo-oligomeric proteins or hetero-oligomeric proteins can be prepared, depending upon the whether the oligomerizing domain(s) of the fusion protein preferentially form hetero-ologimers or homo-oligomers.

Expression Vectors

Recombinant expression vectors for expression of a fusion protein comprising an oligomer-forming domain and a heterologous manunalian protein by recombinant DNA techniques include a DNA sequence comprising a synthetic or cDNA-derived DNA fragment encoding an oligomer-forming domain, linked in frame to a DNA fragment encoding the heterologous protein. These DNA fragments are operably linked to suitable transcription and/or translational regulatory nucleotide sequences, such as those derived from a mammalian, microbial, viral, or insect gene. Examples of regulatory sequences include sequences having a regulatory role in gene expression (e.g., a transcription promoter or enhancer), an operator sequence to control transcription, a sequence encoding an mRNA ribosomal binding site, a polyadenylation site, splice donor and acceptor sites, and appropriate sequences which

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control transcription, translation initiation and termination. In addition, sequences encoding signal peptides can be incorporated into expression vectors. For example, a DNA sequence for a signal peptide (secretory leader) may be operably linked to a DNA encoding a fusion protein comprising an oligomer-forming domain and a heterologous mammalian protein. The signal peptide is expressed as a part of a precursor amino acid sequence; the signal peptide enables improved extracellular secretion of translated fusion polypeptide by a yeast host cell.

Nucleotide sequences are operably linked when the regulatory sequence functionally relates to the DNA encoding the fusion protein. Thus, a promoter nucleotide sequence is operably linked to a DNA encoding a fusion protein if the promoter nucleotide sequence controls the transcription of the DNA encoding the fusion protein. Still further, a ribosome binding site may be operably linked to a sequence for a fusion protein if the ribosome binding site is positioned within the vector to encourage translation.

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Transcription and translational control sequences for mammalian host cell expression vectors may be excised from viral genomes. For example, commonly used mammalian cell promoter sequences and enhancer sequences are derived from Polyoma virus, Adenovirus 2, Simian Virus 40 (SV40), and human cytomegalovirus. DNA sequences derived from the SV40 viral genome, for example, the SV40 origin, early and late promoters, enhancer, splice, and polyadenylation sites may be used to provide the other genetic elements required for expression of a structural gene sequence in a mammalian host cell. Viral early and late promoters are particularly useful because both are easily obtained from a viral genome as a fragment which may also contain a viral origin of replication (Fiers et al., Nature 273:113, 1978). Smaller or larger SV40 fragments may also be used, provided the approximately 250 bp sequence extending from the Hind III site toward the Bgl I site located in the SV40 viral origin of replication site is included.

Exemplary mammalian expression vectors can be constructed as disclosed by Okayama and Berg (Mol. Cell. Biol. 3:280, 1983). A useful high expression vector, PMLSV N1/N4, described by Cosman et al., Nature 312:768, 1984 has been deposited as ATCC 39890. Additional useful mammalian expression vectors are described in EP-A-0367566, and in U.S. Patent Application Serial No. 07/701,415, filed May 16, 1991, incorporated by reference herein. For expression of a type II protein extracellular region, such as OX40-L, a heterologous signal sequence may be added, such as the signal sequence for interleukin-7 (IL-7) described in United States Patent 4,965,195, or the signal sequence for interleukin-2 receptor described in United States Patent Application 06/626,667 filed on July 2, 1984. Another exemplary vector is

pDC406, which includes regulatory sequences derived from SV40, human immunodeficiency virus (HIV), and Epstein-Barr virus (EBV).

Expression vectors transfected into prokaryotic host cells generally comprise one or more phenotypic selectable markers. A phenotypic selectable marker is, for example, a gene encoding a protein that confers antibiotic resistance or that supplies an autotrophic requirement, and an origin of replication recognized by the host to ensure amplification within the host. Other useful expression vectors for prokaryotic host cells include a selectable marker of bacterial origin derived from commercially available plasmids. This selectable marker can comprise genetic elements of the cloning vector pBR322 (ATCC 37017). pBR322 contains genes for ampicillin and tetracycline resistance and thus provides simple means for identifying transformed cells. The pBR322 "backbone" sections are combined with an appropriate promoter and a OX40-L DNA sequence. Other commercially vectors include, for example, pKK223-3 (Pharmacia Fine Chemicals, Uppsala, Sweden) and pGEM1 (Promega Biotec, Madison, WI, USA).

Promoter sequences are commonly used for recombinant prokaryotic host cell expression vectors. Common promoter sequences include β-lactamase (penicillinase), lactose promoter system (Chang et al., Nature 275:615, 1978; and Goeddel et al., Nature 281:544, 1979), tryptophan (trp) promoter system (Goeddel et al., Nucl. Acids Res. 8:4057, 1980; and EP-A-36776) and tac promoter (Maniatis, Molecular Cloning: A Laboratory Manual, Cold Spring Harbor Laboratory, p. 412, 1982). A particularly useful prokaryotic host cell expression system employs a phage λ P_L promoter and a c1857ts thermolabile repressor sequence. Plasmid vectors available from the American Type Culture Collection which incorporate derivatives of the λ P_L promoter include plasmid pHUB2 (resident in E. coli strain JMB9 (ATCC 37092)) and pPLc28 (resident in E. coli RR1 (ATCC 53082)).

Host Cells

Suitable host cells for expression of a fusion protein comprising an oligomerforming domain and a heterologous mammalian protein include prokaryotes and yeast
or higher eukaryotic cells. Prokaryotes include gram negative or gram positive
organisms, for example, E. coli or Bacilli. Suitable prokaryotic host cells for
transformation include, for example, E. coli, Bacillus subtilis, Salmonella
typhlmurium, and various other species within the genera Pseudomonas,
Streptomyces, and Staphylococcus. Higher eukaryotic cells include established cell
lines of mammalian origin. Cell-free translation systems could also be employed to
produce a fusion protein comprising an oligomer-forming domain and a heterologous
mammalian protein using an RNA derived from DNA constructs disclosed herein.

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In a prokaryotic host cell, such as *E. coli*, a fusion protein may include an N-terminal methionine residue to facilitate expression of the recombinant polypeptide in the prokaryotic host cell. The N-terminal Met may be cleaved from the expressed recombinant fusion protein. Prokaryotic host cells may be used for expression of fusion proteins that do not require extensive proteolytic or disulfide processing.

Appropriate cloning and expression vectors for use with bacterial, fungal, yeast, and mammalian cellular hosts are described, for example, in Pouwels et al. Cloning Vectors: A Laboratory Manual, Elsevier, New York, (1985). An expression vector carrying the recombinant fusion protein DNA is transfected or transformed into a substantially homogeneous culture of a suitable host microorganism or mammalian cell line according to methods that are known in the art, to form transfected or transformed host cells that express the fusion protein. Expressed fusion protein will be located within the host cell and/or secreted into culture supernatant fluid, depending upon the nature of the host cell and the gene construct inserted into the host cell.

A fusion protein comprising an oligomer-forming domain and a heterologous mammalian protein may be expressed in yeast host cells, preferably from the Saccharomyces genus (e.g., S. cerevisiae). Other genera of yeast, such as Pichia or Kluyveromyces, may also be employed. Yeast vectors will often contain an origin of replication sequence from a 2µ yeast plasmid, an autonomously replicating sequence (ARS), a promoter region, sequences for polyadenylation, and sequences for transcription termination. Preferably, yeast vectors include an origin of replication sequence and selectable marker. Suitable promoter sequences for yeast vectors include promoters for metallothionein, 3-phosphoglycerate kinase (Hitzeman et al., J. Biol. Chem. 255:2073, 1980) or other glycolytic enzymes (Hess et al., J. Adv. Enzyme Reg. 7:149, 1968; and Holland et al., Biochem. 17:4900, 1978), such as enolase, glyceraldehyde-3-phosphate dehydrogenase, hexokinase, pyruvate decarboxylase, phosphofructokinase, glucose-6-phosphate isomerase, 3-phosphoglycerate mutase, pyruvate kinase, triosephosphate isomerase, phosphoglucose isomerase, and glucokinase. Other suitable vectors and promoters for use in yeast expression are further described in Hitzernan, EPA-73,657.

Yeast vectors can be assembled, for example, using DNA sequences from pBR322 for selection and replication in $E.\ coli$ (Amp^r gene and origin of replication). Other yeast DNA sequences that can be included in a yeast expression construct include a glucose-repressible ADH2 promoter and α -factor secretion leader. The ADH2 promoter has been described by Russell et al. (*J. Biol. Chem. 258:2674, 1982*) and Beier et al. (*Nature 300:724, 1982*). The yeast α -factor leader sequence directs secretion of heterologous polypeptides. The α -factor leader sequence is often inserted between the promoter sequence and the structural gene sequence. See, e.g., Kurjan et

al., Cell 30:933, 1982 and Bitter et al., Proc. Natl. Acad. Sci. USA 81:5330, 1984.
Other leader sequences suitable for facilitating secretion of recombinant polypeptides from yeast hosts are known to those of skill in the art. A leader sequence may be modified near its 3' end to contain one or more restriction sites. This will facilitate fusion of the leader sequence to the structural gene.

Yeast transformation protocols are known to those of skill in the art. One such protocol is described by Hinnen et al., *Proc. Natl. Acad. Sci. USA* 75:1929, 1978. For example, one can select for Trp⁺ transformants in a selective medium, wherein the selective medium consists of 0.67% yeast nitrogen base, 0.5% casamino acids, 2% glucose, 10 µg/ml adenine and 20 µg/ml uracil. Yeast host cells transformed by vectors containing ADH2 promoter sequence may be grown for inducing expression in a "rich" medium. An example of a rich medium is one consisting of 1% yeast extract, 2% peptone, and 1% glucose supplemented with 80 µg/ml adenine and 80 µg/ml uracil. Derepression of the ADH2 promoter occurs when glucose is exhausted from the medium.

Mammalian or insect host cell culture systems could also be employed to express recombinant fusion protein. Examples of suitable mammalian host cell lines include the COS-7 line of monkey kidney cells (ATCC CRL 1651; Gluzman et al., Cell 23:175, 1981), L cells, C127 cells, 3T3 cells (ATCC CRL 163), Chinese hamster ovary (CHO) cells, HeLa cells, BHK (ATCC CRL 10) cell lines, and CV-1/EBNA cells (ATCC CRL 10478). The CV-1/EBNA cell line was derived by transfection of the CV-1 cell line with a gene encoding Epstein-Barr virus nuclear antigen-1 (EBNA-1) and constitutively express EBNA-1 driven from human CMV immediate-early enhancer/promoter. An EBNA-1 gene allows for episomal replication of expression vectors that contain the EBV origin of replication.

Protein Purification

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Purified soluble fusion proteins are prepared by culturing suitable host/vector systems to express the recombinant soluble fusion proteins, which are then purified from culture media or cell extracts, using standard methods of protein purification that are optimized for each individual soluble fusion protein.

For example, supernatants from systems which secrete recombinant protein into culture media are clarified, and concentrated using a commercially available protein concentration filter, for example, an Amicon or Millipore Pellicon ultrafiltration unit.

Following the concentration step, the concentrate can be applied to a suitable purification matrix. Suitable matrices include those useful in affinity chromatography. For example, a suitable affinity matrix can comprise a cognate protein to which the

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fusion proteins binds, or lectin or antibody molecule which binds the fusion protein, bound to a suitable support.

Alternatively, an ion exchange resin can be employed, for example, an anion exchange resin comprising a matrix or substrate having pendant diethylaminoethyl (DEAE) groups, or other suitable anion exchangers. The matrices can be acrylamide, agarose, dextran, cellulose or other types commonly employed in protein purification. Alternatively, a cation exchange step can be employed. Suitable eation exchangers include various insoluble matrices comprising sulfopropyl or carboxymethyl groups.

One or more reversed-phase high performance liquid chromatography (RP-HPLC) steps employing hydrophobic RP-HPLC media, e.g., silica gel having pendant methyl or other aliphatic groups, can be employed to further purify a soluble fusion protein. Size exclusion chromatography will also be useful in purifying soluble fusion proteins. Additionally, hydrophobic supports can also be used under low pressure conditions; an exemplary medium is phenyl-sepharose. Some or all of the foregoing purification steps, in various combinations, can also be employed to provide a homogeneous recombinant protein.

Biological Activity

Biological activity of recombinant, soluble fusion proteins is mediated by binding of the recombinant, soluble fusion protein to a cognate molecule. A cognate molecule is defined as a molecule which binds the recombinant soluble fusion protein in a non-covalent interaction based upon the proper conformation of the recombinant soluble fusion protein and the cognate molecule. For example, for a recombinant soluble fusion protein comprising an extracellular region of a receptor, the cognate molecule comprises a ligand which binds the extracellular region of the receptor. Conversely, for a recombinant soluble fusion protein comprising a ligand, the cognate molecule comprises a receptor (or binding protein) which binds the ligand.

Binding of a recombinant fusion protein to a cognate molecule is a marker for biological activity. Such binding activity may be determined, for example, by competition for binding to the binding domain of the cognate molecule (i.e. competitive binding assays). One configuration of a competitive binding assay for a recombinant soluble fusion protein comprising a ligand uses a radiolabeled, soluble receptor, and intact cells expressing a native form of the ligand. Such an assay is illustrated in Example 4 herein. Similarly, a competitive assay for a recombinant soluble fusion protein comprising a receptor uses a radiolabeled, soluble ligand, and intact cells expressing a native form of the receptor. Instead of intact cells expressing a native form of the cognate molecule, one could substitute purified cognate molecule bound to a solid phase. Competitive binding assays can be performed using standard

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methodology: Qualitative or semi-quantitative results can be obtained by competitive autoradiographic plate binding assays, or fluorescence activated cell sorting, or Scatchard plots may be utilized to generate quantitative results.

Biological activity may also be measured using bioassays that are known in the art, such as a cell proliferation assay. Exemplary bioassays are described in Example 2 herein. The type of cell proliferation assay used will depend upon the recombinant soluble fusion protein. A bioassay for a recombinant soluble fusion protein that in its native form acts upon T cells will utilize purified T cells obtained by methods that are known in the art. Such bioassays include costimulation assays in which the purified T cells are incubated in the presence of the recombinant soluble fusion protein and a suboptimal level of a mitogen such as Con A or PHA. Similarly, purified B cells will be used for a recombinant soluble fusion protein that in its native form acts upon B cells. Other types of cells may also be selected based upon the cell type upon which the native form of the recombinant soluble fusion protein acts. Proliferation is determined by measuring the incorporation of a radiolabeled substance, such as ³H thymidine, according to standard methods.

Yet another type assay for determining biological activity is induction of secretion of secondary molecules. For example, certain proteins induce secretion of cytokines by T cells. T cells are purified and stimulated with a recombinant soluble fusion protein under the conditions required to induce cytokine secretion (for example, in the presence of a comitogen). Induction of cytokine secretion is determined by bioassay, measuring the proliferation of a cytokine dependent cell line. Similarly, induction of immunoglobulin secretion is determined by measuring the amount of immunoglobulin secreted by purified B cells stimulated with a recombinant soluble fusion protein that acts on B cells in its native form, using a quantitative (or semi-quantitative) assay such as an enzyme immunoassay. Example 2 presents such assays.

The relevant disclosures of all references cited herein are specifically incorporated by reference. The following examples are offered by way of illustration, and not by way of limitation.

EXAMPLE 1

This example describes construction of a CD40-L DNA construct to express a soluble CD40-L fusion protein referred to as trimeric CD40-L. CD40-L is a type II transmembrane protein round on activated T cells, that acts as a ligand for the B cell antigen, CD40 (Armitage et al., Nature 357:80, 1992; Spriggs et al., J. Exp. Med. 176:1543, 1992). A gene encoding CD40-L has been cloned and sequenced as described in U.S.S.N. 07/969,703, filed October 23, 1992, the disclosure of which is incorporated by reference herein. CD40-L is a member of the Tumor Necrosis Factor

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(TNF) family of proteins; several members of this family are believed to exist in trimeric form.

Trimeric CD40-L contains a leader sequence, a 33 amino acid sequence referred to as a "leucine zipper" (SEQ ID NO:2), and an eight amino acid hydrophilic sequence described by Hopp et al. (Hopp et al., BioTechnology 6:1204, 1988; SEQ ID NO:1; referred to as Flag®), followed by the extracellular region of human CD40-L (amino acid 50 to amino acid 261 of SEQ ID NOs:3 and 4). The utility of the leader and the Flag® sequences have been described in previously. The 33 amino acid sequence presented in SEQ ID NO:2 trimerizes spontaneously in solution. Fusion proteins comprising this 33 amino acid sequence are thus expected to form trimers or multimers spontaneously.

The construct is prepared by synthesizing oligonucleotides representing a leader sequence, the 33 amino acid sequence described above (SEQ ID NO:2), and the Flag® sequence (SEQ ID NO:1), then ligating the final product to a DNA fragment encoding the extracellular region of human CD40-L (amino acids 50 to 261 of SEQ ID NOs:3 and 4).

The resulting ligation product in expression vector pDC406 was transfected into the monkey kidney cell line CV-1/EBNA (ATCC CRL 10478). The pDC406 plasmid includes regulatory sequences derived from SV40, human immunodeficiency virus (HIV), and Epstein-Barr virus (EBV). The CV-1/EBNA cell line was derived by transfection of the CV-1 cell line with a gene encoding Epstein-Barr virus nuclear antigen-1 (EBNA-1) that constitutively expresses EBNA-1 driven from the human CMV intermediate-early enhancer/promoter. The EBNA-1 gene allows for episomal replication of expression vectors, such as pDC406, that contain the EBV origin of replication.

Once cells expressing the fusion construct are identified, large scale cultures of transfected cells are grown to accumulate supernatant from cells expressing soluble, oligomeric CD40-L. The soluble, oligomeric CD40-L fusion protein in supernatant fluid is purified by affinity purification substantially as described in U.S. Patent 5,011,912. sCD40-L may also be purified using other protein purification methods, as described herein. Silver-stained SDS gels of the soluble, oligomeric CD40-L fusion protein can be prepared to determine purity. Similar methods are used to prepare and purify a trimer-forming construct comprising the extracellular region of murine CD40-L (amino acid 50 to amino acid 260 of SEQ ID NOs:5 and 6). Soluble CD40-L exhibits similar biological activity to that of membrane-bound CD40-L, as shown in Example 2.

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Example 2

This example illustrates B cell proliferative activity and induction of polyclonal immunoglobulin secretion using soluble, oligomeric CD40-L prepared as described in Example 1. Human B cells were purified substantially as described in Armitage et al. (J. Immunol. 150:3671; 1993). Briefly, tonsillar tissue was gently teased and the resulting cell suspension centrifuged over Histopaque® (Sigma, St. Louis, MO). T cell-depleted preparations of cells (E-) were obtained by removing T cells by rosening with 2-aminoethylisothiouronium bromide-treated SRBC (sheep red blood cells) and treatment with B cell Lympho-kwik (One Lambda Inc., Los Angeles, CA) for 1 hour at 37°C to lyse contaminating non-B cells. Peripheral blood mononuclear cells (PBMC) were isolated in the same manner, with the additional step of treating the partially purified cells with 5 mM leucine methyl ester (Leu ME; Sigma, St. Louis, MO) in scrum-free medium for one hour at room temperature prior to the Lympho-kwik step, to remove phagocytic cells.

B cell proliferation was measured with a ³H-thymidine incorporation assay, substantially as described in Armitage et al., supra. Cells were cultured for three days in the presence of soluble, oligomeric CD40-L, alone or in the presence of 5 ng/ml IL-4 (Immunex Corporation, Seattle, WA), 5 µg/ml anti-IgM coated beads (BioRad, Richmond, CA), or a combination of IL-4 and anti-IgM. The results of a representative experiment to evaluate the ability of soluble, oligomeric CD40-L to induce B cell proliferation are shown in Figures 1A and 1B. sCD40-L induced proliferation of tonsillar B cells in the presence of IL-4, anti-IgM, or a combination of these to cofactors (Figure 1A). sCD40-L also induced proliferation of peripheral blood B cells in the presence of IL-4, anti-IgM, or a combination of these to co-factors, and with B cells obtained from some donors, a moderate level of proliferation in the absence of any cofactor (Figure 1B). These results parallel the results obtained with recombinant, membrane-bound CD40-L described in Armitage et al. supra.

Polyclonal immunoglobulin secretion was determined by isotype-specific ELISA on supernatant fluid from 10 day cultures of 1 X 10⁵ B cells per well, substantially as described in Armitage et al., *supra*. Purified B cells were stimulated with a 1:20 dilution of supernatant fluid containing soluble, oligomeric CD40-L (sCD40-L), a 1:20 dilution of control supernatant (control S/N; conditioned medium from cells transfected with vector alone), or transfected CV-1/EBNA cells expressing membrane-bound CD40-L (CV1/CD40L; 3 x 10⁴ cells/well), in the presence or absence of 10 ng/ml of either IL-2, IL-4 (both from Immunex Corporation, Seattle, WA) or IL-10 (Genzyme Corporation, Boston, MA). The results of a representative experiment measuring immunoglobulin secretion are presented in Table 1; values given represent

the quantity of each isotype secreted by the induced B cells in ng/ml, as measured by ELISA.

Table 1: Immunoglobulin Secretion Induced by CD40 Ligand

	ioto 1. attantan		1	CD 40 ENEMIC	
	Medium alone	Medium + IL-2	Medium + IL-4	Medium + IL-10	Isotype
Control S/N	114.9	424.5	69.4	132.2	
sCD40L-3	212.3	2827.5	51.5	1726.5	IgM
CV1/CD40L	91.8	1965.0	97.4	574.1	
Control S/N	16.2	161.0	40.1	22.7	
sCD40L-3	25.8	933.2	122.3	231.9	IgG ₁
CVI/CD40L	2.3	428.0	27.9	247.0	
Control S/N	45.1	44.2	39.6	50.0	
sCD40L-3	56.7	248.0	48.7	353.9	IgA
CVI/CD40L	64.7	513.8	34.7	447.2	
Control S/N	<0.3	<0.3	<0.3	<0.3	
sCD40L-3	<0.3	<0.3	. 67.0	<0.3	· IgE
CV1/CD40L	<0.3	<0.3	77.6	<0.3	

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These results indicated that soluble, oligomeric CD40-L induced polyclonal immunoglobulin secretion in the same manner as membrane-bound CD40-L. IL-2 and IL-10 enhanced secretion of IgM, IgG₁ and IgA; secretion of measurable amounts of IgE occurred only in the presence of IL-4, just as observed for membrane-bound CD40-L. The same pattern of immunoglobulin secretion was present when B cells from several different donors were tested, although the absolute quantities varied from donor to donor. In similar experiments in a murine system, a soluble, oligomeric construct of a murine CD40-L also gave comparable results to membrane-bound murine CD40-L.

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Example 3

This example describes construction of a CD27-L DNA construct to express a soluble, oligomeric CD27-L fusion protein referred to as sCD27L-3. CD27-L is a type II transmembrane protein that binds to the lymphocyte antigen, CD27. CD27 is found on most peripheral blood T cells (Bigler et al., J. Immunol. 141:21, 1988; van Lier et al., Eur. J. Immunol. 18:811, 1988), and a subpopulation of B cells (Maurer et al., Eur. J. Immunol. 20:2679, 1990). CD27-L is a member of the tumor necrosis factor family of cytokines. A gene encoding CD27-L has been cloned and sequenced as

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described in Goodwin et al., Cell 73:447 (1993), and in U.S.S.N. _ , a continuation-in-part of U.S.S.N. 07/941,648, filed September 8, 1992. the disclosures of which are incorporated by reference herein.

The construct encoding sCD27L-3 contains a leader sequence, a 37 amino acid sequence comprising a leucine zipper domain, and the extracellular region of human CD27-L from amino acid 39 to amino acid 193; the nucleoride and amino acids sequences are presented in SEO ID NOs:7 and 7. The construct was prepared by using methods that are well-known in the art to obtain a DNA encoding the extracellular region of CD27-L. Briefly, the extracellular region of CD27-L was amplified from a 10 full-length CD27-L cDNA using a PCR technique. The primers used were derived from the extracellular region of CD27-L (SEQ ID NO:7, nucleotides 222-245, for the 5' primer, and the complement of nucleotides 663-689 for the 3' primer) with addition of sequences encoding desired restriction enzyme sites (ACTAGT, which contains a Spe I site, for the 5' primer, and GCGGCCGC, which contains a Not I site, for the 3' primer). The amplified PCR product, representing the extracellular domain of CD27-L, was cloned into an Spe I/Not I-cut SMAG (pDC206) vector. SMAG vector is a derivative of pDC201 (Sims et al., Science 241:585, 1988) that contains the murine IL-7 leader sequence. The vector was amplified, then cut with Spe I and treated with calf intestinal alkaline phosphatase. Oligonucleotides based on the amino acid sequence of a leucine zipper (SEQ ID NO:1) were synthesized by standard methodology, and ligated with the Spe I-cut vector, to form an expression vector comprising a murine IL-7 leader sequence (Namen et al., Nature 333:571; 1988), a leucine zipper domain, and the extracellular domain of CD27-L. The expression vector was referred to as pDC206/sCD27L-3.

pDC206/sCD27L-3 was co-transfected into the monkey kidney cell line CV-1/EBNA (ATCC CRL 10478) along with a pSV3Neo plasmid. pSV3Neo (Mulligan and Berg, Proc. Natl. Acad. Sci. U.S.A. 78:2072; 1981) is a plasmid which expresses the SV40 T antigen, and thus allows for the episomal replication of the pDC206 plasmid.

Once cells expressing the fusion construct are identified, large scale cultures of transfected cells are grown to accumulate supernatant from cells expressing the soluble. oligomeric CD27-L fusion protein (referred to as sCD27L-3). sCD27L-3 in supernatant fluid is purified by affinity purification substantially as described in U.S. Patent 5,011,912. sCD27L-3 may also be purified using other protein purification 35 methods, as described herein. Silver-stained SDS gels of the soluble, oligometic CD27-L fusion protein can be prepared to determine purity. sCD27L-3 binds to soluble CD27, and inhibits binding of soluble CD27 to cells expressing CD27-L, as described in Example 4.

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Example 4

This example illustrates a binding inhibition activity of sCD27L-3. A soluble form of the human lymphocyte surface antigen CD27 was prepared substantially as described by Fanslow et al., *J. Immunol.* 149:65 (1992), to form a dimeric, Fc fusion construct referred to as CD27.Fc (Goodwin et al., *Cell* 73:447; 1993). CD27.Fc comprises the extracellular region of CD27 and an Fc region from a human IgG₁. sCD27L-3 inhibits binding of CD27.Fc to MP.1 cell, a human, Epstein-Barr virustransformed B cell line that expresses endogenous CD27-L.

Conditioned supernatant fluid from CV-1/EBNA cells transfected with pDC206/sCD27L-3 was titrated in a 96 well plate. A constant amount of CD27.Fc (1 µg/well) was added to each well, followed by 1-2 x 106 MP.1 cells per well, in binding medium (RPMI-1640 containing 1 % bovine scrum albumin, 0.2 % sodium azide and 20 mM HEPES, pH 7.2). The plate was incubated at 37°C for one hour. Cells were washed twice with PBS, then pelleted by centrifugation. ¹²⁵I-mouse anti-human IgG Fc was added to each well at a constant concentration, and the plate incubated for an additional hour at 37°C. The ¹²⁵I-mouse anti-human IgG Fc bound to the CD27.Fc that bound to the MP.1 cells. After the final incubation, cells were harvested over pthalate oil-containing tubes to separate the bound and free ¹²⁵I-mouse anti-human IgG Fc, and the amount of radioactivity quantitated using a gamma counter.

The results of this experiment are presented in Figure 2. sCD27L-3 exhibited a dose-dependent inhibition of the binding of CD27.Fc to MP.1 cells. By comparing the concentration at which the inhibition of binding of CD27.Fc is at 50% to the titration of inhibition by sCD27L-3, it was estimated that the concentration of sCD27L-3 in the conditioned medium was between 18 and 40 μ g/ml. In making this comparison, the MW of sCD27L-3 was estimated to be 135 Kd (estimated MW of extracellular region of CD27-L was 45 Kd, multiplied by three for formation of trimer), and the binding of sCD27L-3 to CD27.Fc was assumed to occur at a molar ratio. The Ki was estimated to be 10 times the Ka, which was 3 x 10⁻⁷M⁻¹, and the initial concentration was assumed to be 1 x 10⁻⁸ M. The results demonstrated that the initial assumption of a concentration of 1 x 10⁻⁸ M was approximately 10-fold too low, and a 1:3 dilution of the supernatant fluid actually gave an estimated concentration of 1 x 10⁻⁷ M.

SEQUENCE LISTING

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(1) GENERAL INFORMATION:
          (i) APPLICANT: Spriggs, Melanie
                       · Srinivasan, Subhashini
         (ii) TITLE OF INVENTION: Mathods of Preparing Soluble.
10
                 Oligomeric Proteins
        (111) NUMBER OF SEQUENCES: 8 -
         (iv) CORRESPONDENCE ADDRESS:
15
               (A) ADDRESSEE: Immunex Corporation
               (B) STREET: 51 University Street
               (C) CITY: Seattle
               (D) STATE: WA
               (E) COUNTRY: USA
20
               (F) ZIP: 98101.
          (v) COMPUTER READABLE FORM:
               (A) MEDIUM TYPE: Floppy disk
               (B) COMPUTER: IBM PC compatible
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               (C) OPERATING SYSTEM: PC-DOS/MS-DOS
               (D) SOFTWARE: PatentIn Release #1.0, Version #1.25
         (vi) CURRENT APPLICATION DATA:
               (A) APPLICATION NUMBER:
30
               (B) FILING DATE:
               (C) CLASSIFICATION:
       (viii) ATTORNEY/AGENT INFORMATION:
               (A) NAME: Perkins, Patricia A
35
               (B) REGISTRATION NUMBER: 34,693
               (C) REFERENCE/DOCKET NUMBER: 1003
         (ix) TELECOMMUNICATION INFORMATION:
               (A) TELEPHONE: (206)587-0430
40
               (B) TELEFAX: (206)233-0644
     (2) INFORMATION FOR SEQ ID NO:1:
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               (A) LENGTH: 8 amino acids
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20	Туг	His 1	le Glu 20	Asn Glu	Ile	Ala	Arg 25	Ile	Lys	Lys	Leu	Ile 30	Gly	Glu	
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/71	INFORMATION	FOR	CEA	TD	MO.A

(1) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 261 amino acids

(B) TYPE: amino acid

(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: protein

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Ile Thr Gln Met Ile Gly Ser Ala Leu Phe Ala Val Tyr Leu His Arg 35 40 45

Arg Leu Asp Lys Ile Glu Asp Glu Arg Asn Leu His Glu Asp Phe Val

Phe Met Lys Thr Ile Gln Arg Cys Asn Thr Gly Glu Arg Ser Leu Ser 25 65 70 75 80

Leu Leu Asn Cys Glu Glu Ile Lys Sar Gln Phe Glu Gly Phe Val Lys 85 90 95

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Met Gln Lys Gly Asp Gln Asn Pro Gln Ile Ala Ala His Val Ile Ser 115 120 125

Glu Ala Ser Ser Lys Thr Thr Ser Val Leu Gln Trp Ala Glu Lys Gly 130 135 140

Tyr Tyr Thr Met Ser Asn Asn Leu Val Thr Leu Glu Asn Gly Lys Gln 40 145 150 155 160

Leu Thr Val Lys Arg Gln Gly Leu Tyr Tyr Ile Tyr Ala Gln Val Thr 165 170 175

45 Phe Cys Ser Asn Arg Glu Ala Ser Ser Gln Ala Pro Phe Ile Ala Ser 180 185 190

Leu Cys Leu Lys Ser Pro Gly Arg Phe Glu Arg Ile Leu Leu Arg Ala 195 200 205

Ala Asn Thr His Ser Ser Ala Lys Pro Cys Gly Gln Gln Ser Ile His 210 215 220

Leu Gly Gly Val Phe Glu Leu Gln Pro Gly Ala Ser Val Phe Val Asn 55 225 230 235 240

Val Thr Asp Pro Ser Gin Val Ser His Gly Thr Gly Phe Thr Ser Phe 245 250 255

60 Gly Leu Leu Lys Leu

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25 ·		(xi)	SE(DENC	Œ DE	SCRI	PTIC	N: 5	SEQ 1	D NO):5:						
																GGA Gly	46
30	1				5					10	5				15	,	
J0																CTT	96
	Leu	Pro	Ala	Ser 20	Het	Lys	Ile	Phe	Het 25	Tyr	Leu	Leu	Thr	Val 30	Phe	Leu	
35				ATG													144
	116	Thr	35 -	Met	116	GTÅ	ser	40	Leu	Phe	ALA	AT	19r 45	Leu	HT2	Arg	
40																GTA	192
40	Arg	50	Asp	Lys	Val	GIU	55	GIU	Val	ASII	Leu	60	GIU	мэр	rne	val	
																TCC	24
45	65	Ile	Lys	Lys	ren	70	Arg	Cys	Asn	rys	75	GIN	GIĀ	Ser	Leu	80 80	
																	28
	Leu	Leu	Asn	Cys	Glu 85	Glu	Met	Arg	Arg	90	Phe	Glu	Хзр	Leu	Val 95	Lys	
50	GAT	RTA	ACG	773	110	222	GAA	GAG	288	***	GAA	110	acc.	TT-T	GNA	ATG	33
				Leu 100	Asn					Lys							33
55	CAA	ÄĞA	GGT	GAT	GAG	GAT	CCT	CAA	ATT	GCA	GCA	CAC	GTT	GTA	AGC	GAA	38
				Asp					Ile								•
	GCC	AAÇ	AGT	AAT	GCA	GCA	TCC	GTT	CTA	CAG	TGG	GCC	AAG	AAA	GGA	TAT	43
60	Ala	Asn 130		Asn	Ala	Ala	Ser 135	Val	Leu	Gln	Trp	Ala 140		Lys	Gly	Tyr	

5	TAT Tyr 145	ACC Thr	ATG Met	AAA Lys	AGC Ser	AAC Asn 150	TTG Leu	GTA Val	ATG Met	CTT Leu	GAA Glu 155	TAA NE A	eta Gee	AAA Lys	CAG Gln	CTG Leu 160	480
-	ACG Thr	GTT Val	AAA Lys	AGA Arg	GAA Glu 165	GGA Gly	CTC Leu	TAT Tyr	TAT Tyr	GTC Val 170	TAC Tyr	ACT Thr	CAA Gln	GTC Val	ACC Thr 175	TTC Phe	528
10											CCA Pro					CTC Leu	576
15	TGG Trp	CTG Leu	AAG Lys 195	CCC Pro	AGC Ser	AGT Ser	GGA Gly	TCT Ser 200	GAG Glu	AGA Arg	ATC Ile	TTA Leu	CTC Leu 205	AAG Lys	GCG Ala	GCA Ala	624
2 0											CAG Gln						672
25		Gly					Gln				TCT Ser 235						720
23	ACT Thr	GAA Glu	GCA Ala	AGC Ser	CAA Gln 245	Val	ATC Ile	CAC His	AGA Arg	GTT Val 250	GGC Gly	TTC Phe	TCA Ser	TCT Ser	TTT Phe 255	G14 GCC	768 -
30			AAA Lys								•-						783
35	(2)	INE	ORMA	TION	FOR	SEQ	ID	NO:6	:								
			(<u>i</u>)	(A (B) LE	ngth Pe:	: 26	o ac	ino [.]		3						
40		,	11)					line									
45		(ĸi)	SEQU	ENCE	DES	CRIP	TION	: SE		NO:						
•	Met 1		Glu	Thr	Tyr 5		Gln	Pro	Ser	Pro 10	Arg	Ser	Val	Ala	Thr 15		
50	Leu	Pro	·Ala	Sez .20		. Lys	Ile	Phe	Met 25		Leu	Leu	Thr	Val 30		Leu	
	Ile	Thr	G1: 39		Ile	G1y	Ser	Val 40		Phe	Ala	Val	Tyr 45		His	Arg	
<i>5</i> 5	Ārg	.Leu 50	•	Lys	Val	. Glu	G1 to 55		Val	. Asn	Leu	His 60		Asp	Phe	Val	
60	Phe 65		Lys	Lys	Lev	1 Lys 70		у Суз	Asn	Lys	Gly 75		Gly	Ser	Leu	Ser 80	

	Leu	Leu	aek	Суз	Glu 85	Glu	Met	Arg	Arg	Gln 90	Phe	Glu	Asp	Leu	Val 95	Lys
5	qeÁ	Ile	Thr	Lou 100	Asn	Lys	Glu	Glu	Lys 105	Lys	Glu	A SD	Ser	Pha 110	Glu	Met
•	Gln	λrg	Gly 115	Asp	Glu	Азр	Pro	Gln 120	Ile	Ala	Ala	His	Val 125	Val	Ser	G1u
10	Ala	Asn 130	Ser	Asn	Ala	Ala	Ser 135	Val	Leu	Gln	Trp	Ala 140	Lys	Lys	Gly	Tyr.
15	Tyr 145	Thr	Met	Lys	Ser	Asn 150	Leu	Va1	Met	Lou	Glu 155	λsn	Gly	Lys	Gln	Leu 160
	Thr	Val	Lys	Arg	Glu 165	Gly	Leu	Tyr	Tyr	Val 170		Thr	Gla	Val	Thr 175	Phe
20	Суз	Ser	Asn	Arg 180	Glu	Pro	Ser	Ser	Gln 185	Arg	Pro	Phe	Ile	Val 190	Gly	Leu
	Trp	Leu	Lys 195	Pro	Ser	Ser	Gly	Ser 200	Glu	Arg	Ile	Leu	Leu 205	Lys	Ala	уја
25	Asn	Thr 210	His	Ser	Ser	Ser	Gln 215	Leu	Cys	Glu	Gln	Gln 220		Val	His	Lau
30	Gly 225	G∫À	Val	Phe	Glu	Leu 230	Gln	Ala	Gly	Ala	Ser 235	Val	Phe	Val	λsn	Val [.] 240
	Thr	Glu	Ala	Ser	Gln 245	Val	Ile	His	Arg	Val 250	Gly	Phe	Ser	Ser	Phe 255	Gly
35	Leu	Leu	Lys	260												
	(2)	INE	ORMA:	rion	FOR	SEQ	ID !	NO:7	:							
40		(i)	(1	A) Li B) T C) S	engti YPE : IRANI	HARAGE BUC	89 ba leic ESS:	aci	pair:	s						
45		(ii)) MO													
		(iii)) RY	POTH	ETIC	AL: 1	10									•
50		(iv) AN	ri-s	ense	: NO				•						
		(vi)	OR:			OURCI ISM:		7 li	gand	tri	mer	(CD2	7L-3	•		_
5 Ş		(ix		A) N	AME/	KEY: ION:		. 686								
60		(ix) FE: () ()	A) N	AME/	KĖY: ION:	sig 39.	_pep	tide	;						

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1	(ix)	FEATURE:	•

(A) NAME/KEY: mat peptide (B) LOCATION: 111.686

(x1) SEQUENCE DESCRIPTION: SEQ ID NO:7:

10	GGAAAAC	TCT (CGAGG	STACE	et as	reco	GGGG	A TC	CCA	H		rc ci he K		al S		53
15	TTT AGA Phe Arg	TAT	ATĆ Ile	TTT Phe -15	GGA Gly	ATT Ile	CCT Pro	CCA Pro	CTG Leu -10	ATC	CTT Leu	GTT Val	CTG Leu	CTG Leu -5	CCT Pro	101
20	GTC ACT Val Thr	AGT Ser	TCT Ser 1	GAC Asp	ÇGT	ATG Met	AAA Lys 5	CAG Gln	ATA Ile	GAG Glu	GAT Asp	AAG Lys 10	ATC Ile	GAA Glu	GAG Glu	149
	ATC CTA Ile Leu 15	Ser														197
25	AAG CTG Lys Leu 30	ATT	GGC Gly	GAG Glu	CGG Arg 35	ACT Thr	AGT Ser	CAG Gln	CGC Arg	TTC Phe 40	GCA Ala	CAG Gln	GCT Ala	CAG Gln	CAG Gln 45	245
30	CAG CTG Gln Leu	CCG Pro	CTC Leu	GAG Glu 50	TCA Ser	CTT	GGG Gly	TGG Trp	GAC Asp 55	GTA Val	Ala GCT	GAG Glu	CTG Leu	CAG Gln 60	CTG Leu	293
35	AST CAC EIH NEA	The	Gly 65	Pro	Gln	Gln	Asp	70	Arg	Leu	Tyr	Trp	Gln 75	GŢĀ	Gly	341
40	CCA GCA Pro Ala	Leu 80	Gly	Arg	Ser	Phe	Leu 85	His	Gly	Pro	Glu	Leu 90	Asp	Lys	Gly	389
	CAG CTA Gln Leu 95	λrg	ATC Ile	CAT His	CGT	GAT Asp 100	GJA GCC	ATC Ile	TAC Tyr	ATG Met	GTA Val 105	CAC His	ATC Ile	CAG Gln	GTG Val	437
45	ACG CTG Thr Leu 110	yj'a ecc	ATC Ile	Cys	TCC Ser 115	TCC	ACG Thr	ACG Thr	AT a	TCC Ser 120	agg Azg	CAC His	CAC His	Pro	ACC Thr 125	485
50	ACC CTG Thr Leu	GCC Ala	GTG Val	GGA Gly 130	ATC Ile	TGC Cys	TCT Ser	CCC Pro	GCC Ala 135	TCC Ser	CGT	AGC Ser	ATC Ile	AGC Ser 140	CTG Leu	533
55 _.	CTG CGT Leu Arg	CTC Leu	AGC Ser 145	TTC Phe	CAC His	CĂA Gln	GCT	TGT Cys 150	ACC Thr	ATT Ile	GTC Val	TCC Ser	CAG Gln 155	CGC	CTG Leu	581
60	ACG CCC Thr Pro	CTG Leu 160	GCC Ala	CGA Arg	GGG Gly	GAC Asp	ACA Thr 165	CTC Leu	TGC Cys	ACC Thr	aac Asd	CTC Leu 170	ACT Thr	gly ggg	ACA Thr	629
•																

Phe Gly Val Gln Trp Val Arg Pro 185 190

								•	29							
Leu	TTG Leu 175	Pro	TCC Ser	CGA	AAC	ACT Thr 180	GAT Asp	GAG Glu	ACC	TTC Pbs	TTT Phe 185	GGA Gly	GTG Val	CAG Gla	TGG	677
		Pro														689,
(2)	INF	DRMA:	TION	FOR	SEQ	ID I	10:8	:								
		(i) :	.(B)		NGTH PE:	210 min	am:	ino :	; ecid:	3						
	(:	Li) I	HOLE	CULE	TYP	E: pi	rote	Ln.								
	(:	KI) :	SEQUI	ENCE	DES	RIP:	rion:	: SE	Q ID	NO:	ð:					
Met -24	Phe	His	Val	Ser -20	Phe	Arg	Tyr	Ile	Phe -15	Gly	Ile	Pro	Pro	Leu -10	Ile	
Leu	Val.	Leu	Leu -5	Pro	Val	Thr	Ser	Ser 1	Asp	Arg	Met	Lys 5	Gln	Ile	Glu	
	10		Glu	•		15					20				-	
25			Ile		30					35					40	
			Gln	45					50					55		
			G1n 60					65					70			
		75					80			•		85			Pro	• •
	90		Lys			95					100					
105			Gln		110					115					120	
			Pro	125					130					135		
		•	Ser 140					145					150			•
Va1	Ser	Gln 155	Arg	Leu	Thr	Pro	Leu 160	λla	Arg	Gly	Азр	Thr 165	Leu	Суз	The	
Asa	Leu 170	Thr	Gly	The	Leu	Leu 175	Pro	Ser	Arg	Asn	Thr 180	λsp	Glu	Thr	Phe	

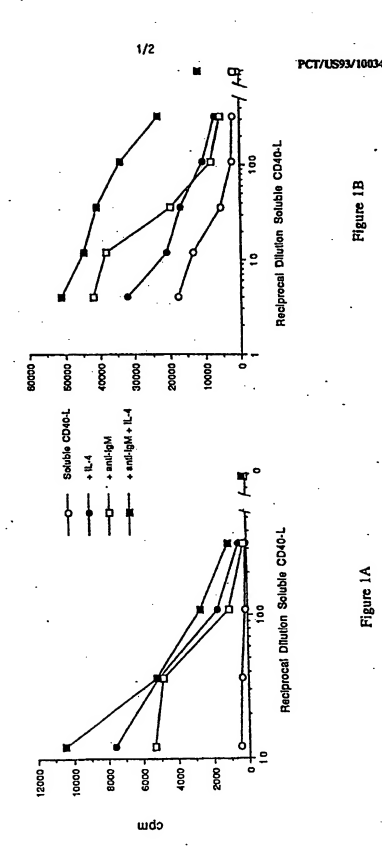
30 CLAIMS

What is claimed is:

- A method of preparing a soluble, oligomeric mammalian protein by culturing a host cell transformed or transfected with an expression vector encoding a fusion protein comprising a leucine zipper domain and a heterologous mammalian protein.
 - 2. The method according to claim 1, wherein the heterologous mammalian protein comprises a transmembrane region of a membrane bound protein.
- 3. The method according to claim 2, wherein the leucine zipper domain comprises a peptide that trimerizes in solution.
 - 4. The method according to claim 3, wherein the leucine zipper domain comprises amino acids 1 through 33 of SEQ ID NO:2.
- 5. The method according to claim 1, further comprising the steps of removing
 the leucine zipper domain from the soluble, oligomeric mammalian protein.
 - 6. The method according to claim 2, further comprising the steps of removing the leucine zipper domain from the soluble, oligomeric mammalian protein.
 - 7. The method according to claim 3, further comprising the steps of removing the leucine zipper domain from the soluble, oligomeric mammalian protein.
- 20 8. The method according to claim 4, further comprising the steps of removing the leucine zipper domain from the soluble, oligomeric mammalian protein.
 - 9. The method according to claim 1, wherein the heterologous mammalian protein comprises a globular mammalian protein.
- 10. The method according to claim 9, wherein the globular mammalian protein25 is a cytokine.
 - 11. A method of preparing a soluble, hetero-oligomeric mammalian protein by culturing a first host cell transformed or transfected with a first expression vector encoding a first fusion protein comprising a first leucine zipper domain and a first heterologous mammalian protein, culturing a second host cell transformed or transfected with a second expression vector encoding a second fusion protein

comprising a second leucine zipper domain and a second heterologous mammalian protein, wherein the first and second leucine zipper domains preferentially form a hetero-oligomer, and combining the first and second fusion proteins under conditions promoting hetero-oligomer formation.

- 12. The method according to claim 11, wherein the first and second heterologous mammalian proteins comprise globular mammalian proteins.
 - 13. The method according to claim 12, wherein the globular mammalian proteins are cytokines.
- 14. The method according to claim 11, wherein the first heterologous
 mammalian protein comprises GM-CSF and the second heterologous mammalian protein comprises IL-3.
 - 15. The method according to claim 11, wherein the first and second heterologous mammalian proteins comprise extracellular regions of transmembrane proteins.
- 15 16. The method according to claim 15, wherein the extracellular regions are from different transmembrane proteins.
 - 17. The method according to claim 16, wherein the extracellular regions are from the same transmembrane protein.
- 18. The method according to claim 1, wherein the fusion protein further comprises a linker sequence.
 - 19. The method according to claim 11, wherein the fusion protein further comprises a linker sequence.



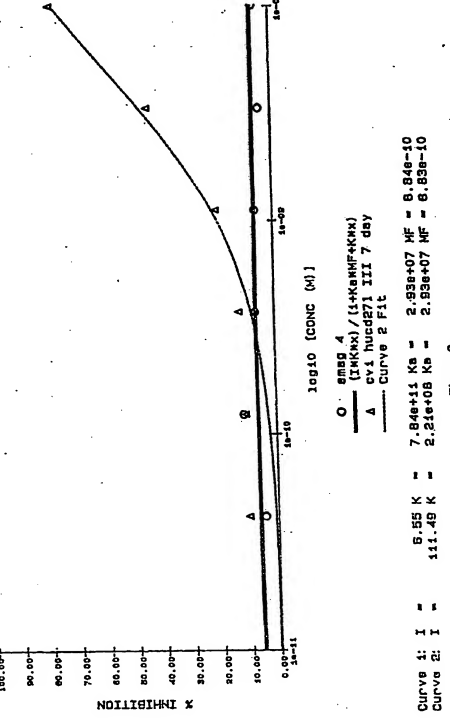


Figure 2

INTERNATIONAL SEARCH REPORT

International application No. PCT/US93/10034

	·			
A. CLASSIFICATION OF SUBJECT MATTER				
PC(5) :C12N 15/12, 15/62 US CL :435/69.7; 536/23.4				
According to International Patent Classification (IPC) or to both national classification and IPC				
B. FIELDS SEARCHED				
Minimum documentation scarched (classification system followed by classification symbols)				
U.S. : 435/69.7, 172.3, 252.3, 320.1; 530/350; 536/23.4				
Documentation searched other than minimum documentation in the extent that such documents are included in the fields searched				
Electronic data base consulted during the interactional search (name of data base and, where practicable, accerd terms used)				
APS, STN/MEDLINE				
search terms: GM-CSF, IL-3, fusions, chimer?, hybrids				
C. DOCUMENTS CONSIDERED TO BE RELEVANT				
Category*	Citation of document, with indication, where a	ppropriate, of the relevant passages Relevant to	chim No.	
			•	
X	The Journal of Immunology, Vol. 148, No. 5, issued 01 March 1.5.9.10			
Y	1992, Kostenly et.al., FORMATION OF A BISPECIFIC 2-4,6-8,11-19			
	ANTIBODY BY THE USE OF LEUCINE ZIPPERS", pages 1547			
	to 1553, see entire document.			
Y	EP, A, 0 325 224 (Sledziewski et.al.) 26 July 1989, see entire 2-4,6-8,11-19			
	document.			
Υ .	US, A, 5,073,627 (Curtis et.al.) 17 December 1991, see entire 2-4,6-8,11-19			
	document.			
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Further documents are listed in the commutation of Box C. See patent family annex.				
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The Accounting published prior to the international files done but help then "A"				
the priority date channel				
12 JAN 31 1994 JAN 31 1994				
Name and mailing address of the ISAUS Authorized officer				
Commissioner of Patents and Trademarks Box PCT		William No		
	L. D.C. 20231	JOHN D. ULM XV-XVIJ 991		
Faceimilie No. NOT APPLICABLE Telephone No. (703) 308-0196				
Form PCT/ISA/210 (second shoot)(July 1992)#				

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